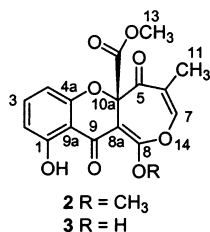


Tim S. Bugni, Valerie S. Bernan, Michael Greenstein, Jeffrey E. Janso, William M. Maiese, Charles L. Mayne, and Chris M. Ireland* Brocaenols A–C: Novel Polyketides from a Marine-Derived *Penicillium brocae*

Page 2014. For compounds **2** and **3**, the R groups are incorrect. As shown below, compound **2** is brocaenol B, and R should equal a methyl. Compound **3** is brocaenol C, and R should equal a hydrogen.



Page 2015. $\Delta\delta^{\text{RS}}$ should be $\Delta\delta^{\text{SR}}$ to denote that values were calculated by subtracting the chemical shift of the R-MTPA ester from the S-MTPA ester. Additionally, the same changes apply to Figure 3 and the text located on the same page. The corrected figure is shown below. Although the symbols were incorrect, the absolute stereochemistry was determined correctly

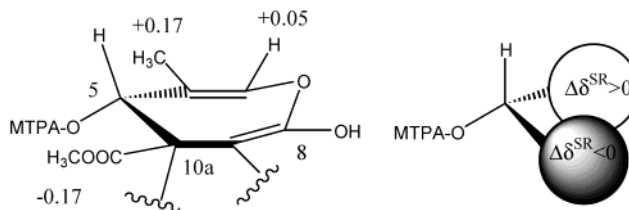


FIGURE 3. Figure 3. Model used to assign the configuration at C-5 with $\Delta\delta^{\text{SR}}$ values in ppm.

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